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(71) Applicant (for all designated States except US): ECLAGEN LIMITED [GB/GB]; Marischal College, Broad Street, Aberdeen AB9 1AS (GB).

(72) Inventors; and

- (75) Inventors/Applicants (for US only): FOTHERGILL, John [GB/GB]; The Granary, Shorehead AB39 2JY, Stonehaven (GB). KEMP, Graham [GB/GB]; Netherton, Sauchen AB59 7JP, Inverurie (GB). BROOKS, Tony [GB/GB]; 10 Sunnybank Place, Aberdeen AB24 3LA (GB). CARR, Frank [GB/GB]; Auris, 23 St. Machar Drive, Aberdeen AB24 3RY (GB).
- (74) Agents: STEBBING, Peter, John, Hunter et al.; Ablett & Stebbing, 45 Lancaster Mews, Lancaster Gate, London W2 3QQ (GB).

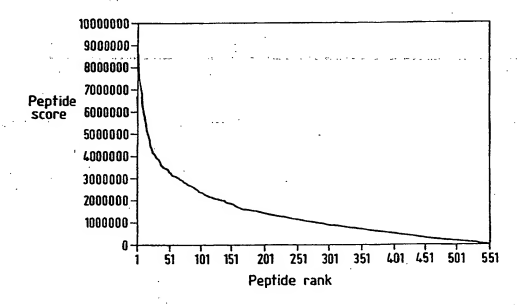
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(57) Abstract

The invention provides a method for the prediction of the binding affinity of a peptide to a major histocompatilibity (MHC) class II molecules comprising; 1) ascertaining the characteristics of a MHC molecule binding groove, 2) presenting a selected peptide to the MHC molecule and ascertaining a first conformation score for each pocket bound peptide side—chain, 3) amending the conformation of each pocket bound peptide side—chain and ascertaining a second conformation score, 4) repeating step 3 with alternative conformations of each peptide pocket bound side—chain, 5) choosing the highest conformation score for each pocket bound peptide side—chain in each binding groove pockets, herein known as "the pocket", and 6) combining the highest conformation score for each pocket and ascertaining a binding score for the complete peptide.

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IDENTIFICATION OF MHC BINDING PEPTIDES

The present invention relates to a new method for the prediction of peptides which bind to major histocompatibility 5 (MHC) class II molecules and to molecules created or modified through the use of these methods.

The immune system of the mammalian organism principally comprises two arms, the cellular immune system and the humoral or antibody-associated immune system. The cellular immune system is centred around the activity of T cells. There are two major classes of T cells, cytotoxic T lymphocytes (CTLs) which attack cells displaying foreign antigen complexed with MHC class I molecules, and helper T cells which react to cells displaying foreign antigens in a complex with MHC class II molecules resulting in the secretion of cytokines which can activate B cells to produce antibody molecules.

Humans express six different MHC class I genes and six 20 different MHC class II genes, which are located on three This leads to considerable allelic highly polymorphic loci. variation in MHC molecules. The MHC class I consist of a α chain and a β_2 -microglobulin, the α -chain is split into three domains α_1 , α_2 and α_3 . α_1 and α_2 form the MHC class I binding 25 groove which contains pockets that bind the side chains and the amino and carboxy termini of any peptide present in the The MHC class II molecules comprise an α -chain and a β -chain, it is the α_1 and β_1 domains which create the MHC class II binding groove. The MHC class II binding groove also 30 contains pockets but it does not bind the end termini of the peptide. For this reason the peptides bound by the MHC class II molecule can be longer and of a more variable length. typical length of peptides complexed with a MHC class I or a MHC class II molecule are 8-10 amino acids and 13-20 amino 35 acids, respectively.

At present only three MHC class II structure are available but

it is believed that the backbone structure of all MHC class II alleles presently identified are similar to that of HLA-DR1. Structures of different alleles can be predicted by using homology modelling. This involves identifying the amino acid differences near the binding groove and using a computer to change the conformation of the side-chains to give favourable steric and electrostatic arrangements and to make the pockets as large as possible. The end result is a three dimensional structure of a MHC class II molecule, which can be used in various experiments.

The ability to predict the peptides in a protein which can bind to a given MHC molecule has great value especially for medical applications. It is known, for example, that in 15 certain auto-immune diseases, T cells react with self-peptides presented by MHC class II molecules. It would be valuable to predict which peptides from auto-immune proteins are presented by MHC class II molecules in these diseases as well as to predict the binding of analogues of these peptides synthesised 20 as potential antagonists for the presentation of In the selection of peptides for synthetic vaccines, the ability to predict MHC class II binding peptides would be advantageous. In addition, where heterologous proteins are developed as medicines or diagnostic imaging 25 agents, it would be advantageous to predict potential MHC class II binding peptides in order to eliminate these from the heterologous proteins before administration to patients.

While studies of peptides complexed with MHC class I molecules
have revealed conserved "anchor" residues at certain positions within the presented peptides, such studies with peptides complexed with MHC class II molecules have been less successful mainly because of the greater length variability of such peptides and the consequent difficulty in aligning their sequences.

Methods for accurately predicting the binding potential of

peptides have been restricted to MHC class I interaction with a peptide. In one method using three-dimensional structures of MHC class I molecules, peptide binding is ranked in ascending order according to the energy values determined.

5 This method requires that the MHC structure be known, or that there is an obvious molecular model for the MHC structure. An identical method is said to be available for MHC class II but it does not consider the longer average length of the peptide and the open-ended peptide binding groove of MHC class II molecules. Neither does it use the best potential conformation of peptide amino acid side-chains and, therefore the binding energies calculated are only approximations.

Another drawback of using the same method for MHC class I and
15 MHC class II peptide binding is that the binding of peptides
to MHC class II is less dependant on strict allele-specific
binding motifs than peptides binding to MHC class I.
Individual amino acids in the peptide play a more significant
role in MHC class II binding than MHC class I such that the
20 conformation of amino acid side-chains is proportionally more
important to the accuracy of binding analysis. Therefore,
known methods do not provide a general method for analysing
the binding of peptides to three-dimensional structures of MHC
class II. There is thus a need for improved methods for
25 predicting the MHC class II binding potential of peptides.

An object of this invention is to provide a method for accurately predicting the binding affinity of a peptide fragment binding to a MHC class II molecule.

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Another object of this invention is to provide a computer conditioned to perform the task of predicting the binding affinity of a peptide fragment binding to a MHC class II molecule.

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A yet further object of this invention is to provide a vaccine derived from the peptide fragment whose binding affinity to

MHC class II molecules has been determined.

Another object of this invention is to provide a pharmaceutical composition which comprises a peptide whose binding affinity to MHC class II molecules has been determined.

According to the first aspect of this invention, there is provided a method for the prediction of the binding affinity of a peptide and a major histocompatibility (MHC) class II molecules comprising;

- 1) ascertaining the characteristics of a MHC molecule binding groove,
- 2) presenting a selected peptide to the MHC molecule and 15 ascertaining a first conformation score for each pocket bound peptide side-chain,
 - 3) amending the conformation of each pocket bound peptide side-chain and ascertaining a second conformation score,
- 4) repeating step 3 with alternative conformations of each 20 peptide pocket bound side-chain,
 - 5) choosing the highest conformation score for each pocket bound peptide side-chain,
- 6) combining the highest conformation score for each pocketbound peptide side-chain and then ascertaining a binding score25 for the peptide.

It is particularly desirable to then compile information on all peptide fragments in a protein and compare the binding scores. It is preferable if the conformation of the backbone of the peptide fragment is also altered and the conformation score and the binding score is then reassessed.

The method of this invention thus involves assessing a binding score for all possible candidate peptides by considering the predicted three-dimensional conformations and interactions between the MHC and the peptide in the complex. The computed score indicates the predicted binding affinity for the

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particular peptide binding with the MHC allele and can be used to predict whether the peptides are likely to bind, or not.

Preferably, the conformation score for each pocket bound 5 peptide side-chain is ascertained by considering at least one of the following parameters:

- a) the steric overlap between the pocket bound peptide residue bound in the pocket and an atom forming the pocket; this is value B,
- 10 b) the number of hydrogen bonds which can be formed between the pocket bound peptide residue and an atom forming the pocket; this is value C,
 - c) the strength of electrostatic interactions between any polar atoms of the pocket bound peptide residue and any polar
- 15 atoms forming the pocket; this is value D, and
 - d) the number of favourable contacts between the pocket bound peptide residue and the MHC residues forming one of the pockets; this is value E.
- The conformation score for each peptide is computed based upon the predicted atomic interactions between each of the pocket bound peptide residues and MHC pockets. The geometric constraints imposed on the peptide by the shape of the MHC binding groove play an important part of the scoring function.
- Favourable packing arrangements between peptide and MHC sidechains are rewarded by the scoring function, whilst
 arrangements involving steric overlap are penalised.
 Alternative conformation are tried for MHC residues if an MHC residue overlaps with a peptide side chain.

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If no preferable conformation can be found the MHC side-chain is returned to its original conformation. In the event of more than a pocket residue side-chain overlapping with a pocket bound peptide side chain, the pocket residue side chains are adjusted in order of overlap severity, with the pocket residue side-chain which has the most severe overlap being adjusted first.

In preferred embodiments the steric overlap between the pocket bound peptide residue and the atoms forming the pocket can not be greater than 0.35 Angstroms, otherwise the residue is deemed unable to fit in the pocket.

5

Conveniently a favourable contact occurs when an atom from an MHC residue and an atom from the peptide residue have their centres separated by no more than the sum of their radii plus 0.5 Angstroms and are not overlapping.

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Preferably the values B to E are imported into a first equation to give a conformation score(Z). The first equation is $Z_n = (cK_2C) - (cK_3D) + (cK_4E) - (cK_1B)$, where cK_1 to cK_4 are constants and n is the number of the pocket.

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The value of cK_1 is between 50 and 150. Preferably between 75 and 125.

The value of cK_2 is between 1000 and 2000. Preferably between 20 1250 and 1750.

The value of cK_3 is between 250 and 750. Preferably between 350 and 650.

25 The value of cK_4 is between 500 and 1500. Preferably between 750 and 1250.

Conveniently the Z_n value for a pocket is multiplied by a coefficient, L, depending on the pockets importance in binding, to give a second Z_n value. The value L is in the range of 0.001 to 5. Larger pockets are considered more important in determining which peptide can bind, compared with the other smaller pockets, so the scores contributed by each pocket are weighted in proportion to the amount of the peptide side-chain buried by the surface of the MHC molecule. When binding to MHC class II molecules, peptides have shown high similarity in the degree to which their side-chains are buried

by the MHC surface, despite having dissimilar sequences.

Preferably all the Z_n values are summed to give a value J. Value J is the overall contributing score of all the pockets for a certain conformation of the peptide fragment.

Conveniently the MHC residue is paired with the pocket-bound peptide residue if an atom from the MHC residue and an atom from the pocket-bound peptide residue have their centres separated by no more than the sum of their van der Waal radii plus one Angstrom.

In a preferred embodiment a value A_n is calculated by summing the pairwise interaction frequencies of paired residues. As 15 for the Z_n value, preferably the value A_n for a pocket is multiplied by a coefficient, X, depending on the pockets importance in binding. Preferably X is between 0.001 and 5.

Conveniently the A_n value for the pockets are summed to give 20 a value P.

In a preferred embodiment the binding score is ascertained by at least one of the following parameters

- a) the number of groove-bound hydrophobic residues; this is25 value F,
 - b) the number of non groove-bound hydrophilic residues; this is value G,
 - c) the number of peptide residues deemed to fit within their respective binding pocket; this is value H.

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Preferably values F, G, H, J and P are imported into a second equation to give a first binding score, Y.

Conveniently the second equation is $Y=J*F^2*(G*H+1)+P$.

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However, in the alternative, the term He, which evaluates the hydrophobicity of the pocket bound peptide side chains using

a hydrophobicity scale disclosed in Janin et al [1979] Nature, 277 pg 491, can also be used to determine the Y value. Accordingly, $Y=(bK_2C)-(bK_3D)+(bK_4E)-(bK_1B)+(bK_5He)+P$. The scale used in Janin et al to measure hydrophobicity has a range from 5 -1.8 for lysine to 0.9 for cysteine.

Ιt is known that peptides having favourable · hydrophobic/hydrophobic interactions with solvent and MHC atoms have a higher binding affinity. Accordingly, it is 10 preferable to include the term He.

The value of bK_1 is between 1 and 10. Preferably between 1 and 5.

15 The value of bK_2 is between 20 and 60. Preferably between 30 and 50.

The value of bK₃ is between 300 and 900. Preferably between 450 and 750.

The value of bK_4 is between 1 and 20. Preferably between 5 and 15.

The value of bK_5 is in between 1 and 800. Conveniently 25 between 100 and 600. Preferably between 100 and 400.

In a preferred embodiment determination of the conformation score and the binding score are repeated for each pocket and each conformation of the peptide residue in said pocket. 30 conformation of the peptide is altered by rotating a side chain of the peptide residue by a pre-determined amount. this way all possible conformations of the peptide side-chain in the pocket can be studied and the best or most likely conformation can be chosen to obtain the binding score.

The conformation of the backbone of the peptide fragment is changed by modelling the conformation of the backbone on any

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one of 167 backbones which have been previously generated, based on human and murine crystallographic structures of MHC class II peptide complexes. The backbone conformation and the conformation of the peptide fragment side chains are altered systematically until the conformation score and the binding score of every possible conformation has been determined.

Conveniently the steps are repeated using different peptides from a protein.

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In preferred embodiments the binding scores (Y) for different peptides are tabulated and compared. Peptides with the highest scores are predicted to have the highest binding affinity for the particular MHC allele.

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In a preferred embodiment the method of determining the binding affinity of a peptide residue for an MHC class II molecule is used in the manufacture of a vaccine derived from a peptide identified by said method.

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Preferably the method of determining the binding affinity of a peptide residue for an MHC class II molecule is used to remove potentially immunogenic sequences from a protein and thus reduce said proteins immunogenicity when administered to 25 an organism.

Using the afore-detailed method it is possible to predict the peptides from an auto-immune protein which are presented by MHC class II molecules. Thereafter, it is possible to synthesise peptides which would be antagonists to the presentation of such peptides by the MHC class II molecules. It is also possible to determine any proteins in a vaccine containing heterologous proteins which might result in the stimulation of T cells due to their presentation on MHC class II molecules. These proteins could then be altered or removed depending on their function in the vaccine.

According to a second aspect of the invention there is provided a computer conditioned to receive information characterising a peptide bound to the MHC molecule and to utilise said information to perform a procedure having the 5 following steps;

- 1) ascertaining the characteristics of a MHC molecule binding groove;
- 2) presenting a selected peptide, which is selected by a predetermined program, to the MHC molecule and ascertaining10 a first conformation score:
 - 3) amending the conformation of the peptide, by way of a predetermined program, and ascertaining a second conformation score;
 - 4) repeating step 3 with other conformations of the peptide;
- 15 5) selecting the peptide conformation with the highest conformation score; and
 - 6) calculating the binding score from the conformation score.

Preferably the above detailed procedure also includes a step 20 (7) which comprises repeating steps 1-4 with other peptide fragments in the protein to generate information on all peptide fragments in a protein so that a comparison can be made of the strength of the binding between the peptide and the MHC molecule.

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Conveniently the above detailed procedure further comprising a step (8) which comprises altering the conformation of the backbone of the peptide fragment.

The use of a computer in such a task is important because there are hundreds of calculations to perform per peptide fragment. A computer conditioned to perform the task can systematically change the conformation of the side chains and the backbone of the peptide fragment while calculating the conformation score and the binding score.

According to a third aspect of the invention there is provided

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a pharmaceutical composition made by determining the binding affinity of a peptide for a MHC class II molecule.

A pharmaceutical composition is thus engineered to contain a peptide which is presented by an MHC class II molecule and which therefore stimulates the bodies cellular immune system. Alternatively the pharmaceutical composition is engineered so that it does not include peptides which significantly stimulate the immune system.

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The invention will now be described, by way of illustration only, with reference to the following examples, tables and figures accompanying the specification.

Figure 1 shows a graphical representation of the binding score distribution of all 554 13-mer Influenza haemagglutinin peptides bound to HLA-DRB1*0101.

Figure 2 shows a graphical representation of the binding score distribution of all 554 13-mer Influenza haemagglutinin peptides bound to HLA-DRB1*0401.

Table 1 shows the value for all the factors required to determine the binding score for the 15 peptides from Influenza haemagglutinin which have the highest binding affinity for HLA-DRB1*0101.

Table 2 shows the value for all the factors required to determine the binding score for the 15 peptides from Influenza 30 haemagglutinin which have the highest binding affinity for HLA-DRB1*0401.

Table 3 lists the sequence difference between HLA-DRB1*0101 and HLA-DRB1*0401.

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Table 4 shows the torsion angles of the mutated side chains in HLA-DRB1*0401.

Example 1

The following method was used to confirm that the peptide PKYVKQNTLKLAT, has a high affinity binding for the MHC molecule HLA-DRB1*0101.

- 5 The conformation score was calculated as follows for an oligomeric peptide having thirteen amino acid residues, herein known as a 13-mer peptide:
- a) Calculate the steric overlap between the pocket bound 10 peptide residue in the binding groove and an atom forming the pocket; this is value B.
- b) Count the number of hydrogen bonds which could be formed between the pocket bound peptide residue and atoms forming the
 pocket; this is value C.
 - c) Calculate the strength of electrostatic interactions between any polar atoms of the pocket bound peptide residue and any polar atoms forming the pocket; this is value D.
 - d) Count the number of favourable contacts between the pocket bound peptide residue and atoms forming the pocket; this is value E.
- 25 These values were then transformed into a conformation score (Z) by using the following equation:

$$Z_n = (cK_2C) - (cK_3D) + (cK_4E) - (cK_1B)$$

where cK_1 to cK_4 are constants and n is the number of the 30 pocket. CK_1 , cK_2 , cK_3 and cK_4 are equal to 100, 1500, 500 and 1000 respectively.

The conformation of each rotatable side chain of the pocket bound peptide bound residue was then altered by 30° and the conformation score was recalculated.

The above steps were repeated for each of the pockets and the

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highest conformation score for each of the pockets was used to determine the binding score.

The binding score was determined by establishing values for 5 the following parameters:

- a) the number of groove-bound hydrophobic residues; this is value F.
- b) the number of non groove-bound hydrophilic residues; this is value G.
- 10 c) the number of peptide residues deemed to fit within their respective binding groove; this is value H.

The conformational scores for pockets one and five were doubled and then all the conformational scores were summed to give a value J.

The above values were then imported in to the following equation in order to determine the binding score:

 $J*F^2*(G*H+1)+P$

The binding scores for all the 13-mer peptides from Influenza Haemagglutinin binding with MHC molecule HLA-DRB1*0401 were calculated and the resultant top 15 binding scores are presented in Table 1. PKYVKQNTLKLAT has the 8th highest binding affinity for HLA-DRB1*0101 from all 554 possible overlapping 13-mer peptides.

Table 1

	Rank	Seq.	Peptide	Binding	P	В	С	D	E	F	G	н
				Score			1		1			
	1	328	NTLKLATGMRNVP	9382500	15012	0.00	1		27	4	6	5
5	2	453	IDLTDSEMNKLFE	8288922	17964	0.72	1		40	3	6	5
	3	373	NSEGTGQAADLKS	7520420	10661	0.68	0	+0.01	30		7	
	4	504	HDVYRDEALNNRF	7211042	15527	0.56	1	-0.05	31	3	6	5
	5	119	PDYASLRSLVASS	7174962	17351	0.68	1		40	4	4	5
	6	461	NKLFEKTRRQLRE	7049469	19407	0.79	0	+0.01	56	2	7	5
10	7	122	ASLRSLVASSGTL	6922064	16346	0.09	0		25	4	4	5
	8	322	PKYVKQNTLKLAT	6765975	18217	1.82	1		56	3	5	5
	9	458	SEMNKLFEKTRRQ	6156822	16617	0.30	4	+0.08	44	2	7	5
	10	513	NNRFQIKGVELKS	6096900	14052	1.32	3	-0.01	30	4	7	4
	11	439 -	YNAELLVALENQH	5890199	14198	0.60	1		33	4	4	5
15	12	63	STGKICNNPHRIL	5887908	12776	0.75	5	-0.05	31	3	6	5
	13	50	IEVTNATELVQSS	5503551	14297	0.95	2	+0.06	39	3	5	5
	14	262	NSNGNLIAPRGYF	5284475	10102	0.09	1		21	4	5	5
Į	15	257	DVLVINSNGNLIA	5239292	17028	1.35	2		35	3	4	5

20

Example 2

A method as described in Example 1 was used to confirm that the peptide PDYASLRSLVASS from Influenza haemagglutinin, has high affinity binding for the MHC molecule HLA-DRB1*0401.

The structure of HLA-DRB1*0401 is not known but a three dimensional model was constructed based on the known structure of HLA-DRB1*0101 by homology modelling. 10 amino acid differences between the two molecules were identified (see Table 2) and HLA-DRB1*0101 was mutated using the molecular modelling package 'Quanta' to produce a model of HLA-DRB1*0401.

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Then the side-chain conformations of the 10 amino acids were adjusted interactively. In most cases, torsion angles were chosen which resulted in little or no steric overlap between the mutated residues and surrounding atoms. In the case of 5 non-conserved residues which were either charged or whose side-chains were able to form hydrogen bonds, the potential to form favourable interactions was also considered. placement of 13H, 28D and 71K was such that these residues were able to form a favourable electrostatic arrangement 10 whilst at the same time, having minimum steric overlap with surrounding atoms. In the case of 30Y, this residue was positioned such that its hydroxyl group was situated close to the side-chain of 9E, where a hydrogen bond may be formed. The torsion angles chosen for the 10 mutated amino acid 15 residues were calculated in accordance with the standard conventions and are listed in Table 3.

The binding scores for all 13-mer peptides from Influenza Haemagglutinin binding with MHC molecule HLA-DRB1*0401 were calculated and the resultant top 15 binding scores are presented in Table 4. PDYASLRSLVASS has the 9th highest binding affinity for HLA-DRB1*0401 from all 554 possible overlapping 13-mer peptides.

Table 2

Seq. Pos.	HLA-DRB1*0101	HLA-DRB1*0401
.b9	Tryptophan	Glutamic acid
b11	Leucine	Valine
b13	Phenylalanine	Histidine
b26	Leucine	Phenylalanine
b28	Glutamic acid	Aspartic Acid
b30	Cysteine	Tyrosine
b31	Isoleucine	Phenylalanine
b33	Asparagine	Histidine
b37	Serine	Tyrosine
b71	Arginine	Lysine
	b11 b13 b26 b28 b30 b31 b33 b37	b9 Tryptophan b11 Leucine b13 Phenylalanine b26 Leucine b28 Glutamic acid b30 Cysteine b31 Isoleucine b33 Asparagine b37 Serine

Table 3

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	Residue	C1	C2	c 3	C4
	b9	-61°	-71°	-2°	
•	b11	168°			
	b13	-38°	-63°		
20	b26	170°	57°		
	b28	-174°	-15°		
	p30	-174°	41°		
	b31	-119°	-13°		
	b33	-95°	-2°		
25	b37	-116°	-2°		
	b71	−97°	-45°	172°	9°

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Table 4

			,,									
	Rank	Seq.	Peptide	Binding	P	В	С	D	E	F	G	н
				Score								
	1	453	IDLTDSEMNKLFE	3070823	6559	0.36	0		42	3	6	5
	2	3,73	NSEGTGQAADLKS	2988447	4182	0.36	0	+0.01	32	4	7	5
5	3	328	NTLKL:ATGMRNVP	2899375	4639	0.00	1		27	4	6	5
 -	4	122	ASLRSLVASSGTL	2894599	6819	0.03	0		24	4	4	5
	5	72	HRILDGIDCTLID	2820446	4623	0.60	1	+0.16	28	4	6	5
	6	461	NKLFEKTRRQLRE	2662369	7203	0.36	0	-0.11	50	2	7	5
-	7	119	PDYASLRSLVASS	2616648	6184	0.11	1		32	4	4	5
10	8	188	DNFDKLYIWGIHH	2615259	5429	0.58	0	·	29	5	6	4
·	9	322	PKYVKQNTLKLAT	2515861	6407	0.46	2		44	3	5	5
	10	232	NIGSRPWVRGLSS	2488137	4818	0.41	0	-0.02	35	4	5	5
	11	504	HDVYRDEALNNRF	2353661	4965	0.05	1	-0.07	25	3	6	5
	12	135	EFITEGFTWTGVT	2208179	3543	0.07	1		20	4	5	5
15	13	251	TIVKPGDVLVINS	2176819	5259	0.10	0		16	5	5	4
	14	257	DVLVINSNGNLIA	2107570	6673	0.71	2 [.]		40	3	4	5
_ [15	439	YNAELLVALENQH	2035430	4795	0.03	1	·	26	4	4	5

20 Example 3

A library of backbones were constructed by examining the crystal structure of the HLA-DR1 complexed with SEB superantigen. This results in a collection of homogenous peptides within the MHC binding groove. The atomic positions of the peptide backbone, as shown in the PDB file produced from the crystal, were considered to be the `representative' backbone conformation of a peptide which binds to HLA-DR1.

30 Each of the peptide backbone conformations from the known MHC class II crystallographic structures are taken and after being transformed to the same frame of reference as the 'representative' peptide had the differences between their $C\alpha/C\beta$ positions and those of the 'representative' peptide

calculated. These differences summarise the variability of $C\alpha/C\beta$ atomic positions between the known peptides and the representative peptide.

5 The differences were doubled to take into account the fact that the variability of peptides thus far crystallised may not fully represent the true variability of peptides binding to MHC class II molecules. The differences were then used to define regions within which peptide $C\alpha$ and $C\beta$ atoms centres are constrained to lie.

An exhaustive search was then made through candidate peptide backbones. Starting from the 'representative' peptide candidates are generated by adjusting backbone ϕ and ψ angles in ten degree steps from the N-terminus to the C-terminus. An adjustment was rejected if it led to any $C\alpha$ or $C\beta$ atom centre being outside the allowed region, derived above. An adjustment which did not violate the constraint results in a new backbone conformation which is stored within the peptide backbone library.

The x, y, and z co-ordinates of atoms in the backbones designated 0, 14, 62, 65, 75, 93, 104, 107, 112, 118, 129, 134, 141, 144 are given in Tables 5 to 18.

Table 5

Backbone 0				
Atom Number	Atom type	Position in peptide	х у	Z
0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41	N A C O C N A C C	0000011111222223333344444555556666677777788	19.913	22.078 22.516 23.352 22.593 22.044 22.536 21.770 20.547 22.743 22.510

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Table 5 continued

	Atom	Atom	Position	x		
	Number			*	У	Z
	Mamper	type	in peptide			
	42	С	. 8	-4.839	75.618	20.504
5	43	0	8.	-4.505	74.687	21.236
	44	СВ	8	-3.924	75.908	18.149
	45	N	9	-6.093	76.041	20.436
	46	CA	9	-7.113	75.382	21.236
	47	С	9	-7.976	74.424	20.403
	48	0	9	-8.366	74.742	
	49	CB	9	- 7.963	76.413	21.973
	,50	N	10	-8.203	73.232	20.971
10	51	CA	10	-8.995	72.149	20.365
	52	С	10	-10.492	72.527	20.200
	53	0	10	-10.962	73.563	20.702
	54	CB	10	-8.830	70.835	21.191
	55	N	11	-11.238	71.661	19.523
	56	CA	11	-12.654	71.907	19.270
	57	С	11	-13.603	71.483	20.395
	58	0	11	-13.661	70.302	20.800
15	59	CB	11	-13.072	71.269	17.940
	60	N	12	-14.360	72.481	20.852
	61	CA	- 12	-15.363	72.337	21.898
ĺ	62	С	12	-14.758	72.166	23.281
	63	0	12	-14.785	71.069	23.853
	64	CB	12	-16.320	71.168	21.577

Table 6

	Backbone 14					
	Atom	Atom	Position	·x	У	z
5	Number	type	in peptide		. -	
10		type N CA C O CB N CA		0.000 18.281 16.799 16.250 0.000 16.174 14.768 14.098 13.053 14.090 14.723 14.182 12.659 11.952 14.470 12.242 10.845 10.219 10.898 10.669 8.980 8.245	0.000 86.637 86.756 87.880 0.000 85.601 85.553 84.393 84.588 86.846 83.223 82.013 82.164 82.431 80.825 82.022 82.086 80.681 79.694 82.621 80.660 79.430	0.000 22.405 22.715 22.720 0.000 22.931 23.287 22.569 21.908 22.869 22.680 22.093 21.901 22.884 22.994 20.649 20.317 20.423 20.101 18.906 20.898 21.010
25	21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41 42 43 44	CA COCNA COC	4 4 4 5 5 5 5 5 6 6 6 6 6 7 7 7 7 7 8 8 8 8 8 8	6.863 6.283 8.071 6.427 5.135 4.084 4.171 5.174 3.174 2.100 1.349 1.703 1.139 0.381 -0.441 -1.906 -2.505 -0.346 -2.392 -3.758 -4.704 -4.316 -4.043	79.586 80.680 79.059 78.504 78.479 77.942 76.770 77.593 78.832 78.470 77.248 76.776 79.635 76.781 75.677 76.139 76.533 74.551 76.454 75.537 74.404 76.313	20.344 20.413 22.472 19.710 19.082 20.074 20.468 17.848 20.452 21.336 20.769 19.678 21.492 21.550 21.137

Table 6 continued

Atom Number	Atom type	Position in peptide	х	У	z
45 46 47 48 49 50 51 52 53 54 55 57 58 59 61 62 63 64	N CA C O CB N CA C O CB N CA C O CB	9 9 9 9 10 10 10 10 11 11 11 11 11 12 12 12 12	-5.873 -6.881 -7.500 -7.243 -7.964 -8.250 -8.934 -10.393 -11.075 -8.914 -10.781 -12.127 -13.058 -13.254 -12.180 -13.551 -14.474 0.000 18.356 0.000	76.084 75.338 74.285 74.336 76.275 73.372 72.354 72.786 73.192 71.043 72.710 73.032 71.846 70.984 73.341 71.844 70.830 -12.127 0.000 0.000	20.610 21.313 20.371 19.159 21.818 20.978 20.229 19.976 20.928 20.996 18.708 18.320 18.640 17.770 16.834 19.872 20.305 73.032 -12.127 0.000

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Table 7

Backbone 62					
Atom Number	Atom type	Position in peptide	х	У	z
0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 1 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41 41 41 41 41 41 41 41 41 41 41 41 41	N CA C O CB N CA C	00000111112222233333444445555666667777788888	0.000 18.315 16.796 16.173 0.000 16.231 14.791 14.286 13.659 14.135 14.144 11.890 14.518 12.614 11.890 14.518 10.289 11.105 10.289 11.105 10.596 8.328 6.861 6.157 8.377 6.490 5.140 4.171 4.543 5.006 21.975 1.074 0.052 -0.931 -2.553 -0.941 -3.658 -4.518 -5.155	0.000 86.971 86.979 87.867 0.000 85.979 85.876 84.665 84.823 83.487 82.241 82.249 82.071 82.071 82.071 82.071 82.591 80.623 79.356 78.440 78.440 78.440 78.740 77.740 78.7	0.000 22.396 22.404 21.780 0.000 23.075 23.216 22.451 21.380 22.652 22.989 22.404 22.212 23.195 23.305 20.629 20.734 20.783 19.218 20.800 20.852 20.395 20.876 22.268 19.470 18.978 20.139 21.055 17.909 20.577 19.503 21.418 21.102 21.418 21.102 21.661 22.174 20.754



Table 7 continued

Atom Number	Atom type	Position in peptide	x	У	Z
44 45 46 47 48 49 50 51 52 53 55 55 57 59 61 62 64	CB N C C O C N C C O C N C C O C B C C O C N C C O C D C C O C D C C O C D C C O C D C C O C D C C O C D C C O C D C C O C D C C O C D C D	8 9 9 9 9 10 10 10 10 11 11 11 11 12 12 12 12	-4.845 -6.623 -7.650 -8.161 -8.197 -8.802 -8.492 -9.030 -10.518 -11.258 -8.887 -10.869 -12.232 -13.047 -13.155 -12.284 -14.366 0.000 18.332 0.000	76.793 76.163 75.345 74.329 74.658 76.215 73.143 72.107 72.390 72.730 70.758 72.271 72.455 71.182 70.312 72.752 71.124 70.022 -12.232 0.000 0.000	18.545 21.113 21.696 20.655 19.460 22.170 21.153 20.315 20.029 20.964 21.000 18.754 18.336 18.641 17.764 16.847 19.871 20.291 72.455 -12.232 0.000

Table 8

Backbone 65	5		·	• *	
Atom Number	Atom type	Position in peptide	х	У	z
0 123456789011231456171892122342567899312333453673894142	N A C O C N A C C	00000111112222233333444445555566666777778888	0.000 18.487 16.990 16.510 0.000 16.279 14.844 14.178 13.234 14.301 14.699 14.144 12.616 11.950 14.457 12.150 10.206 10.895 10.491 9.029 8.376 6.930 6.309 8.365 6.484 5.139 4.150 4.487 4.985 3.002 1.959 0.861 0.752 1.360 0.134 -0.959 -1.631 -3.087 -4.156 -5.496	0.000 86.641 86.870 87.999 0.000 85.796 85.866 84.664 84.830 87.132 83.484 82.241 82.381 82.065 80.624 79.773 82.819 79.322 80.419 79.322 80.419 79.322 80.419 79.322 80.419 77.274 78.731 78.731 78.731 78.731 77.634 77.533 77.634 77.533 77.634 77.533 77.634 77.533 77.634 77.533 77.634 77.533 77.634 77.533 77.634 77.533 77.634 77.533 77.634 77.533 77.634 77.533 77.634 77.6	0.000 22.418 22.533 22.287 0.000 22.868 23.065 22.417 21.612 22.424 22.746 22.248 22.089 23.038 23.212 20.895 20.608 20.484 19.902 19.314 21.065 20.993 20.491 20.801 22.364 19.718 19.212 20.363 21.280 18.142 20.275 21.246 20.665 19.433 21.628 21.573 21.187 20.366 20.039 22.422 20.048 19.326 19.676

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Table 8 continued

	Atom Number	Atom type	Position in peptide	х у	z
5	43 44 45 46 47 48 49 50	O CB N CA C O CB N	8 8 9 9 9 9 9 10	-6.146 75.692 -3.906 76.820 -5.817 76.283 -7.058 75.736 -7.606 74.721 -7.311 74.855 -8.071 76.849 +8.339 73.746	18.775 17.831 20.964 21.439 20.416 19.219 21.649
10	51 52	CA C	10 10	-8.959 72.751 -10.421 73.147	19.824
	53 54 55	O CB N	10 10 11	-10.685 73.773 -8.919 71.398 -11.294 72.734	20.799 20.735
	56 57 58 59	CA C O CB	11 11 11 11	-12.689 73.067 -13.474 71.860 -13.031 71.253 -12.873 74.262	20.085 19.099
15	60 61 62 63	и СА С	12 12 12 12 12	-14.572 71.556 -15.436 70.486 0.000 -12.689	20.766 20.348 73.067
	64	СВ	12	18.675 0.000 0.000 0.000	

Table 9

Backbone 75						
Atom Number	Atom type	Position in peptide	×	У	Z	
0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 38 39 40 40 40 40 40 40 40 40 40 40 40 40 40	N C C O C N C C	00000111112222233333344445555566666777778	0.000 18.442 16.947 16.452 0.000 16.265 14.823 14.466 14.197 14.218 14.505 14.144 12.615 11.895 14.601 10.808 10.331 11.176 10.592 9.013 8.414 6.322 8.478 6.482 5.116 4.181 4.609 4.932 2.974 1.974 0.736 0.349 1.576 0.206 -0.980 -1.844 -1.778 -2.952	0.000 86.539 86.419 86.839 0.000 85.822 85.676 84.417 84.487 86.875 83.290 82.013 81.942 81.727 80.882 82.159 82.078 80.615 79.709 82.592 80.465 79.160 79.245 80.304 78.609 77.470 76.823 77.969 77.470 76.823 77.969 77.867 79.855 76.879 77.867 77.879 77.8	0.000 22.377 22.136 21.066 0.000 23.109 23.048 22.277 21.057 22.338 22.985 22.404 22.214 23.200 23.308 20.971 20.626 20.726 20.772 19.213 20.789 20.836 20.377 20.544 22.251 19.793 19.354 20.577 21.629 18.483 20.389 21.420 20.910 19.748 21.788 21.478 20.470 20.071 22.745 20.088	

Table 9 continued

Atom Number	Atom type	Position in peptide	х	У	Z
42 43 44 45 46 47 48 49 50 51 52 53 54 55 57 59 60 61 62 63 64	C O CB N CC O CB N CC O CB CC O CB	8 8 8 9 9 9 9 10 10 10 10 11 11 11 11 11 12 12 12 12	-5.324 -6.195 -3.604 -5.491 -6.786 -7.424 -7.209 -7.681 -8.142 -8.840 -10.312 -10.616 -8.772 -11.149 -12.546 -13.321 -12.815 -12.741 -14.483 -15.343 0.000 18.817 0.000	76.483 76.435 76.435 76.194 75.859 74.747 74.729 77.087 73.864 72.797 73.196 73.833 71.532 72.774 73.108 72.011 71.509 74.445 71.674 70.702 -12.546 0.000 0.000	19.579 18.698 17.762 20.865 21.391 20.535 19.314 21.388 21.219 20.556 20.334 19.314 21.394 21.275 21.233 20.475 19.460 20.540 21.023 20.406 73.108 -12.546 0.000

.Table 10

Backbone 93						
Atom	Atom	Position	x y	7	z	
Number	type	in peptide				
0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 31 33 33 33 33 33 33 33 33 33 33 33 33	N C C O C N C C	0000011111222223333334444455556666677777788888	0.000 18.249 16.910 16.646 0.000 14.782 14.078 12.999 13.932 14.712 14.144 12.613 11.912 14.484 12.179 10.775 10.163 10.712 10.564 9.085 8.374 7.026 6.568 8.130 6.482 5.203 4.087 4.298 5.163 2.980 1.833 1.164 1.603 0.839 0.169 -0.585 -2.092 -2.667 -0.300 -2.639 -4.853 -4.314	0.000 86.312 86.341 87.271 0.000 85.351 85.213 83.978 84.095 86.434 82.828 81.568 81.568 81.689 81.688 80.454 79.206 78.295 78.295 77.229 78.741 78.572 77.213 76.513 79.695 76.838 74.729 75.687 76.338 74.729 75.344 74.368	0.000 21.629 22.345 23.139 0.000 22.027 22.662 22.127 21.505 22.357 22.345 21.812 22.828 22.959 20.587 20.300 20.176 19.439 19.005 20.925 20.882 20.159 20.925 20.882 20.036 21.936 20.936 20.935 20.066 20.991 17.954 19.876 20.726 20.434 19.510 20.486 21.254 21.080 21.037 22.086 21.254 21.080 21.037 22.086 21.254 21.080 21.037 22.086 21.254 21.080 21.037 22.086 21.254 21.080 21.254 21.080 21.254 21.080 21.254 21.080 21.254 21.2	

Table 10 continued

Atom Number	Atom type	Position in peptide	×	У	z
44 45 46 47 48 49 50 51 53 54 55 57 58 59 61 62 63 64	CB N CA C O CB N CA C O CB N CA C O CB CB	8 9 9 9 9 10 10 10 10 11 11 11 11 11 12 12 12 12	-4.445 -6.082 -6.974 -8.018 -8.754 -7.679 -8.002 -8.947 -10.274 -10.348 -9.194 -11.256 -12.539 -13.542 -13.224 -12.418 -14.678 -15.731 0.000 18.616 0.000	75.782 75.791 75.097 74.312 74.928 76.089 72.999 72.137 72.891 73.727 70.899 72.533 73.179 72.288 71.836 74.524 72.054 71.281 -12.539 0.000 0.000	18.223 20.882 21.769 20.948 20.163 22.679 21.144 20.488 20.269 19.356 21.332 21.087 21.038 20.278 19.167 20.343 20.925 20.326 73.179 -12.539 0.000

Table 11

Backbone 10	4				
Atom Number	Atom type	Position in peptide	x	У	z
0 1 2 3 4 5 6 7 8 9 10 11 2 13 14 15 16 17 18 19 20 12 22 32 24 25 26 27 28 29 30 31 33 33 34 35 36 37 38 38 39 40 40 40 40 40 40 40 40 40 40 40 40 40	N A C O C N A C O C N A C O C N A C O C N A C O C N A C O C N C C O C N A C O C N C C	000001111122222333333444455555666667777788888	0.000 18.400 16.914 16.453 0.009 14.763 14.059 14.693 14.693 14.125 14.465 12.594 11.945 14.465 12.104 10.699 10.406 8.902 8.250 6.415 9.001 4.164 5.130 4.011 4.164 5.968 1.823 1.166 1.718 0.047 -0.707 -2.793 -0.435 -4.974 -4.444	0.000 86.585 86.850 87.991 0.000 85.793 85.897 84.662 84.778 87.122 83.511 82.241 82.372 82.807 81.169 82.048 80.604 79.315 82.801 80.444 79.315 79.315 79.315 76.935 77.091 78.680 77.138 77.091 78.680 77.138 77.1	0.000 22.355 22.523 22.296 0.000 22.880 23.128 22.593 21.971 22.421 22.810 22.404 22.277 23.241 23.424 21.093 20.837 20.723 20.317 19.548 21.120 21.029 20.290 20.160 22.420 19.817 19.147 20.165 20.975 18.066 20.975 18.066 20.975 18.066 20.975 18.066 20.947 20.656 19.864 20.708 21.334 21.135 21.083 22.129 22.267 19.873 19.670 20.684 21.228

Table 11 continued

Atom Number	Atom type	Position in peptide	х у г
44 45 46 47 48 49 50 51 52 53 54 55 56 57 58 59 60 61 62 63 64	CB N CA C O CB N CA C O CB N CA C O CB	8 9 9 9 9 10 10 10 11 11 11 11 11 12 12 12 12	-4.550 75.803 18.256 -6.200 75.824 20.911 -7.100 75.134 21.794 -8.146 74.358 20.969 -8.997 74.991 20.328 -7.800 76.129 22.704 -8.007 73.038 21.000 -8.934 72.175 20.320 -10.266 72.919 20.092 -10.341 73.752 19.177 -9.181 70.924 21.145 -11.249 72.557 20.907 -12.537 73.194 20.850 -13.514 72.294 20.086 -13.514 72.297 18.847 -12.421 74.537 20.152 -14.310 71.549 20.860 -15.320 70.695 20.297 0.000 -12.537 73.194 18.422 0.000 -12.537 0.000 0.000 0.000

Table 12

Backbone 107						
Atom Number	Atom type	Position in peptide	х	Y	Z .	
0 123456789 10112314 156789 1012234256789 2031333356373894014243	N A C C O C N A C	0000011111222223333344445555666667777788888	0.000 18.468 16.971 16.491 0.000 16.260 14.825 14.159 13.215 14.282 14.680 14.125 12.597 11.931 14.438 12.131 10.723 10.187 10.876 10.472 9.010 8.357 6.911 6.290 8.346 6.465 5.120 4.131 4.469 4.966 2.983 1.940 0.842 0.733 1.341 0.115 -0.978 -2.002 -1.726 -1.650 -3.106 -4.175 -5.514 -6.165	0.000 86.641 86.870 87.999 0.000 85.796 85.866 84.664 84.830 87.132 83.484 82.241 82.381 82.065 82.065 80.624 79.773 82.818 80.419 79.322 80.350 78.340 79.322 80.350 78.340 77.274 78.731 78.547 77.634 77.533 76.952 76.952 76.921 76.242 75.692	0.000 22.418 22.533 22.287 0.000 22.868 23.065 22.417 21.612 22.424 22.746 22.248 22.089 23.038 23.212 20.895 20.608 20.484 19.902 19.314 21.065 20.993 20.491 20.801 22.364 19.718 19.212 20.363 21.280 18.142 20.275 21.246 20.665 19.433 21.628 21.573 21.187 20.366 20.039 22.422 20.048 19.326 19.326 19.326 19.326	



Table 12 continued

Atom Number	Atom type	Position in peptide	· x	Υ.	z
44 45 46 47 48 49 50 51 52 53 54 55 57 58 60 61 62 63 64	CB N CA C O CB N CC O CB N CC O CB N CC O CB	8 9 9 9 10 10 10 10 11 11 11 11 12 12 12 12	-3.925 -5.836 -7.077 -7.625 -7.330 -8.090 -8.358 -8.977 -10.440 -10.703 -8.938 -11.313 -12.708 -13.493 -13.050 -12.892 -14.591 -15.455 0.000 18.675 0.000	76.283 75.736 74.721 74.855 76.849 73.746 72.751 73.147 73.773 71.398 72.734 73.067 71.860 71.253 74.262 71.556 70.486 -12.708	17.831 20.964 21.439 20.416 19.219 21.649 20.940 20.108 19.824 18.787 20.799 20.735 20.635 20.635 20.085 19.099 19.715 20.766 20.348 73.067 -12.708 0.000

Table 13

12		
Atom Position	х у	z
type in peptide	_	
Atom Position	0.000 0.000 18.408 86.726 16.919 86.606 16.449 87.028 0.000 0.000 16.215 86.005 14.774 85.858 14.438 84.649 14.190 84.795 14.176 87.097 14.470 83.480 14.125 82.241 12.600 82.176 11.849 82.152 14.572 81.057 12.224 82.187 10.839 82.083 10.319 80.669 11.133 79.744 10.674 82.359 9.001 80.583 8.361 79.323 6.868 79.411 6.126 80.158 8.500 78.961 6.516 78.676 5.150 78.615 4.229 78.301 4.706 77.734 4.995 77.540 2.976 78.716 1.986 78.455 0.948 77.449 1.060 77.031 1.291 79.747 0.020 77.088 -1.045 76.194 -2.219 76.999 -2.062 78.205 -1.517 75.422 -3.314 76.286	2 0.000 22.399 22.121 21.041 0.000 23.077 22.981 22.125 20.907 22.337 22.761 22.093 21.872 22.858 22.932 20.598 20.557 20.692 18.745 20.701 20.960 20.585 21.239 22.429 19.537 19.095 20.291 21.285 18.033 20.149 21.158 20.621 19.459 21.158 20.540 20.540 20.540 20.301 22.353 20.301

Table 13 continued

Atom Number	Atom type	Position in peptide	×	У	z
46 47 48 49 50 51 52 53 54 55 57 58 59 61 62 63 64	CA C O CB N CA C O CB N CA C O CB C O CB	9 9 9 10 10 10 10 11 11 11 11 12 12 12 12	-7.676 -7.858 -7.297 -8.883 -8.598 -8.898 -10.415 -11.204 -8.455 -10.740 -12.112 -12.689 -12.384 -12.211 -13.459 -14.109 0.000 18.708 0.000	75.631 74.446 74.482 76.549 73.451 72.298 72.236 72.400 71.034 72.040 71.910 70.583 69.523 71.942 70.705 69.563 -12.112 0.000 0.000	21.417 20.447 19.341 21.338 20.920 20.116 19.842 20.784 20.832 18.569 18.163 18.695 18.128 16.648 19.770 20.354 71.910 -12.112 0.000

Table 14

Backbone 11	8				
Atom Number	Atom type	Position in peptide	×	У .	Z
0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 27 28 29 30 31 32 33 33 34 35 36 37 38 39 40 41 42 42 43 44 44 45 46 46 47 47 47 47 47 47 47 47 47 47 47 47 47	NACOOC	00000111112222233333344444455555666667777778888888	0.000 18.471 16.968 16.498 0.0046 14.795 14.271 13.620 14.318 14.591 14.592 10.692 10.	0.000 86.536 87.742 0.000 85.669 84.435 86.904 82.045 82.045 82.045 82.067 82.0625 82.0625 82.0625 82.0625 82.0625 82.0625 82.0625 82.0625 82.0625 82.0625 82.0625 82.0625 82.0625 82.0625 83.2625 83.	0.000 22.407 22.266 21.755 0.000 22.686 22.663 21.986 20.922 21.884 22.589 22.093 21.934 22.957 20.366 20.479 20.343 18.958 20.756 20.479 20.343 18.958 20.756 20.171 20.070 22.301 19.106 20.177 20.866 19.106 19.106 20.177 20.866 19.202 21.203 21.2

Table 14 continued

Atom Number	Atom type	Position in peptide	х	У	Z
45 46 47 48 49 51 55 55 57 58 59 61 62 64	N CA C O CB N CA C O CB N CA C O CB N CA C O CB	9 9 9 9 10 10 10 10 11 11 11 11 12 12 12 12	-6.612 -7.652 -8.169 -8.200 -8.795 -8.513 -9.059 -10.544 -11.281 -8.931 -10.894 -12.254 -13.135 -13.091 -12.328 -13.856 -14.763 0.000 18.754 0.000	76.081 75.273 74.268 74.604 76.156 73.083 72.056 72.355 72.703 70.703 72.239 71.287 70.187 72.490 71.586 70.632 -12.254 0.000 0.000	

Table 15

Backbone 12	9				
Atom	Atom	Position	x	У	z
Number	type	in peptide	-	1	2
0 1234567891011 121314151617181920 212232425262728930 313233435637383940 4124344454647	N C C O C N C C	0 0 0 0 0 0 1 1 1 1 1 2 2 2 2 2 2 2 3 3 3 3 3 3 4 4 4 4 4 4 5 5 5 5 5 5 5 5	0.000 18.495 17.099 16.668 0.000 16.409 15.079 14.331 14.767 14.125 12.611 11.911 14.358 12.194 10.652 9.165 10.655 8.445 7.047 6.608 8.305 6.442 4.079 4.373 1.658 1.658 1.658 1.658 1.658 1.658 1.658 1.658 1.658 1.658 1.6594 1	0.000 86.291 86.364 87.449 0.000 85.228 85.125 83.972 84.204 86.412 82.758 81.927 80.407 81.901 82.082 80.727 80.085 83.058 80.348 79.131 79.462 80.330 78.588 77.714 78.866 78.588 77.714 78.866 79.690 76.690 76.044 74.657 75.226 75.414 74.533 75.774 75.079 74.330	0.000 22.091 22.686 23.137 0.000 22.645 23.217 22.570 21.766 22.900 22.404 22.245 23.367 20.297 19.349 19.522 21.074 20.297 19.349 19.522 21.074 20.376 22.102 19.647 19.113 20.180 20.993 17.881 20.145 21.044 20.993 17.881 20.145 21.044 20.630 19.673 21.018 21.0

- 42 -

Table 16 continued

Atom Number	Atom type	Position in peptide	×	У	Z
5 47 48 49 50 51 52 53 54 55 56 57 58 59 60 61 62 63 64	C O CB N CA C O CB	9 9 10 10 10 10 11 11 11 11 11 12 12 12 12	-8.036 -8.773 -7.698 -8.021 -8.966 -10.293 -10.367 -9.213 -11.275 -12.558 -13.561 -13.243 -12.437 -14.696 -15.750 0.000 18.616 0.000	74.312 74.928 76.089 72.999 72.137 72.891 73.727 70.899 72.533 73.179 72.288 71.836 74.524 72.054 71.281 -12.558 0.000 0.000	20.948 20.163 22.679 21.144 20.488 20.269 19.356 21.332 21.087 21.038 20.278 19.167 20.343 20.925 20.326 73.179 -12.558 0.000

Table 17

Atom Atom Position x y Number type in peptide 0 0.000 0.000 1 CA 0 18.454 86.485 2 C 0 16.950 86.573 3 0 0 16.481 87.224 4 CB 0 0.000 10.020 85.918 11.47.776 85.918 0 13.601 84.	
Number type in peptide 0 N 0 0.000 0.000 1 CA 0 18.454 86.485 2 C 0 16.950 86.573 3 O 0 16.481 87.224 4 CB O 0.000 0.000 5 N 1 16.227 85.893 6 CA 1 14.776 85.918 7 C 1 14.776 85.918 7 C 1 14.252 84.663 8 O 1 13.601 84.752 9 CB 1 14.299 87.132 10 N 2 14.573 83.520 11 CA 2 14.106 82.241 12 C 2 12.572 82.273 13 O 2 11.868 82.483 14 CB 2 14.499	0.000 22.460 22.266 21.305 0.000 23.151
1 CA 0 18.454 86.485 2 C 0 0 16.950 86.573 3 O 0 0 16.481 87.224 4 CB 0 0.000 0.000 5 N 1 16.227 85.893 6 CA 1 14.776 85.918 7 C 1 14.252 84.663 8 O 1 13.601 84.752 9 CB 1 14.299 87.132 10 N 2 14.573 83.520 11 CA 2 14.106 82.241 12 C 2 12.572 82.273 13 O 2 11.868 82.483 14 CB 2 14.499 81.135 15 N 3 12.141 82.099 16 CA 3 10.736 82.054 17 C 3 10.224 80.605 18 O 3 11.035 79.698 19 CB 3 10.489 82.573 20 N 4 8.911 80.468 21 CA 4 8.289 79.172	22.460 22.266 21.305 0.000 23.151
22 C 4 6.823 79.286 23 O 4 6.108 80.179 24 CB 4 8.338 78.611 25 N 5 6.465 78.404 26 CA 5 5.118 78.352 27 C 5 4.147 78.042 28 O 5 4.521 77.295 29 CB 5 4.999 77.280 30 N 6 2.972 78.656 31 CA 6 1.943 78.430 32 C 6 1.943 78.430 33 O 6 1.265 76.719 34 CB 6 1.130 79.697 35 N 7 0.034 76.991 36 CA 7 -0.938 75.983 37 C 7 -2.338 76.622 38 O 7 -2.577 77.649 40 N 8 -3.173	22.452 21.387 22.349 23.055 22.559 22.400 23.398 23.523 21.156 20.855 20.973 21.214 19.449 20.833 20.868 20.405 20.882 22.279 19.478 18.981 20.138 21.054 17.911 20.055 21.033 20.562 19.488 21.234 21.401 21.081 20.985 21.637 22.150 20.156 19.995 20.641

Table 17 continued

Atom Number	Atom type	Position in peptide	×	У	Z
48 49 50 51 52 53 54 55 57 58 59 60 61 62 63 64	O CB N CA C O CB N CA C O CB C O CB	9 10 10 10 10 10 11 11 11 11 11 12 12 12 12	-6.531 -9.013 -8.822 -8.965 -10.460 -11.065 -8.334 -10.983 -12.353 -12.732 -12.400 -12.548 -13.373 -13.836 0.000 18.541 0.000	73.205 75.766 73.200 71.925 71.616 70.945 70.836 72.148 71.910 70.452 69.551 72.168 70.294 69.000 -12.353 0.000 0.000	20.765 21.470 20.803 20.155 19.939 20.788 21.005 18.840 18.476 18.805 18.020 16.992 19.958 20.380 71.910 -12.353 0.000

Table 18

Backbone 14	4			· · · · ·	
Atom	Atom	Position	x	У	7
Number	type	in peptide		2	Z
0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 22 22 23 24 25 26 27 28 29 30 31 33 33 33 34 35 36 36 36 36 36 36 36 36 36 36 36 36 36	N A C C O C N A C C O C N A C C O C N A C C O C N A C C O C N A C C O C N A C C O C N A C C O C C N C C O C C N C C O C N C C	00000111111222223333334444455555666667777778888888	0.000 18.480 16.967 16.431 0.000 16.308 14.861 14.262 13.512 14.341 14.630 14.106 12.568 14.581 10.094 10.578 10.098 10.1088 10.1	0.000 86.428 86.551 87.361 0.000 85.727 84.643 84.919 87.091 82.241 82.287 80.628 79.120 80.628 79.228 79.228 79.335 78.598 79.228 79.335 77.643 77.643 77.643 77.643 77.948 77.9	0.000 22.392 22.343 21.553 0.000 23.153 23.256 22.416 21.454 22.745 22.767 22.093 22.092 23.158 22.796 20.899 20.743 20.667 20.273 19.479 21.077 21.020 20.292 20.167 22.424 19.822 19.162 20.737 18.081 20.765 19.676 21.481 21.553 21.512 21.027 21.512 22.174 20.198 20.198 21.512 21.512 21.512 21.512 21.512 22.174 20.843 20.931 18.722

Table 18 continued

Atom Number	Atom type	Position in peptide	х	У	Z
45 46 47 48 49 50 51 52 53 54 55 56 57 58 60 61 62 63	N CA C O CB N C C O CB N C C O CB N C C O CB	9 9 9 9 10 10 10 10 11 11 11 11 12 12 12 12	-6.623 -7.669 -8.201 -8.407 -8.801 -8.360 -8.894 -10.383 -11.124 -8.745 -10.734 -12.097 -12.907 -12.859 -12.150 -13.575 -14.414 0.000 18.465 0.000	76.144 75.348 74.343 74.731 76.243 73.106 72.067 72.344 72.681 70.719 72.224 72.403 71.126 70.178 72.700 71.155 70.059 -12.097 0.000 0.000	21.290 21.873 20.832 19.672 22.347 21.286 20.448 20.162 21.097 21.133 18.886 18.469 18.774 17.977 16.980 19.921 20.322 72.403 -12.097 0.000

Example 4

The following method was used to identify high affinity binding peptides from Myelin Basic Protein (MBP). The binding affinities for a set of MBP peptides to HLA-DRB1*0401 have been experimentally determined and published. This set includes all possible 13 amino acid peptides from the MBP sequence which have a hydrophobic anchor residue at the P3 position. It is known that only such peptides bind to HLA-DR molecules with detectable affinity.

The same homology model of HLA-DRB1*0401 was used for this example as was used in Examples 1 and 2.

- 15 For each of the 13-mer peptides from the experimental determined set, a binding score was calculated as follows:
- a) Calculate the steric overlap between the pocket bound peptide residue in the binding groove and an atom forming the
 20 pocket; this is value B.
 - b) Count the number of hydrogen bonds which could be formed between the pocket bound peptide residue and atoms forming the pocket; this is value C.
 - c) Calculate the strength of electrostatic interactions between any polar atoms of the pocket bound peptide residue and any polar atoms forming the pocket; this is value D.
- 30 d) Count the number of favourable contacts between the pocket bound peptide residue and atoms forming the pocket; this is value E.
- e) These values were then transformed into a conformation 35 score (Z) by using the following equation:

 $Z_n = cK_2C - cK_3D + cK_4E - cK_1B$

Where K_1 to K_4 are constants and n is the sequence position of the peptide residue (numbered from 1 to the N-terminus to 13 at the C-terminus). K_1 , K_2 , K_3 and K_4 are equal to 100, 1500, 500 and 1000, respectively.

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The conformation of each rotatable side-chain of the peptide residue was then altered by 15 degrees and the conformation score was recalculated.

10 The above steps were repeated for each residue of the peptide and the highest conformation score for each peptide residue was sued to determine the conformation score for the peptide.

At the point, the entire proceedings for establishing the conformation score for the peptide were repeated another 166 times, each time using a different peptide backbone form the library of peptide backbones.

The combination of peptide backbone and peptide side-chain conformations which gave the best conformation was then used to determine a binding score for the peptide.

The binding score was determined by establishing values of the following parameters:

- a) Calculate the steric overlap between the pocket bound peptide residue in the binding groove and an atom forming the pocket; this is value B.
- 30 b) Count the number of hydrogen bonds which could be formed between the pocket bound peptide residue and atoms forming the pocket; this is value C.
- c) Calculate the strength of electrostatic interactions 35 between any polar atoms of the pocket bound peptide residue and any polar atoms forming the pocket; this is value D.

- d) Count the number of favourable contacts between the pocket bound peptide residue and atoms forming the pocket; this is value E.
- 5 e) Calculate the hydrophobicity of the pocket bound peptide side chains using a hydrophobicity scale disclosed in Janin et al.
- f) Calculate the number of MHC pocket residues which are paired with the pocket bound peptide residues. Pairing takes place if the centre of an atom from the MHC pocket residue and the centre of an atom from the pocket bound peptide residues are no more than the sum of their van der wall radii plus one Angstrom. The value An is calculated by summing the number of paired residues, where n is the number of the pocket. The values of An taking into account the pockets importance in binding are summed to give a value P.

The above values were then imported in to the following 20 equation in order to determine the binding score (Y):

$Y=P+bK_2C-bK_3D+bK_4E-bK_1B+bK_5He$

Wherein the values bK_1 , bK_2 , bK_3 , bK_4 and bK_5 are 2, 40, 600, 25 10 and 200 respectively.

As can be seen from the results in Table 19 the top four predicted scores pertain to four peptides which appear within the top five best binders.

Table 19

BB	PEPTIDE AF	FINITY	BINDING	D	E	F	В	P	НС
			SCORE						
104	HFFKNIVTPRTPP	40 .	4729	-0.12	11	17	97.7	3580	1 4
107	VHFFKNIVTPRTP	135	2125	-0.19	12	15	284.5	2255	1.5 0.2
104	PVVHFFKNIVTPR	161	4528	-0.06	15	12	337.6	4565	1.4
104	FSWGAEGQRPGF G	298	5205	-0.15	12	10	169.7	4670	-0.2
104	KGFKGVDAQGTLS	480	4353	-0.09	9	13	68.2	3145	
112	KYLATASTMDHAR	479	2672	-0.09	13	15		1480	1.9
129	SKYLATASTMDHA	601	498	-0.08	11	13	275.7	620	2.4 0.4
141	RGLSLSRFSWGAE	1213	4140	-0.05	17	16	81.4	3455	
62	TGILDSIGRFFGG	2942	337	0.04	21	17	25.3	-8	1.7
0	RFFGGDRGAPKRG	3403	3218	-0.24	20	14	369.1	3100	-0.6
104	NIVTPRTPPPSQG	6615	1971	0	10	11	305	2090	1.6
14	DSIGRFFGGDRGA	7268	1904	-0.08	8	15	37.3	16 4 0	8.0
0	SRFSWGAEGQRPG	8352	1735	-0.08	20	13	466.8		0.2
104	SKIFKLGGRDSRS	8494	1387	-0.1	10	10	149.2	1965	8.0
118	SDYKSAHKGFKGV	8510	1884	-0.27	14	14	14.2	825	2.8
65	STMDHARHGFLPR	8860	1886	-0.21	14	15		775.	0.7
104	NPVVHFFKNIVTP	12870	1347	-0.11	12	10	191.3 332.5	1410	2.2
104	GTLSKIFKLGGRD	16000	4162	-0.11	17	10	118	1690	0.2
93	GRFFGGDRGAPKR	18467	244	-0.11	8	9	· · · -	3775	1.1
75	KIFKLGGRDSRSG	25358	2185	-0.13	19	12	161	-175	2.3
0	FGYGGRASDYKSA	26397	1301	-0.12	15	15	279.4	2060	1.4
0	PGFGYGGRASDYK	35200	3485	0.01	14	13	306.1	1530	-0.4
144	GILDSIGRFFGGD	44400		-0.09	21	1.4	183.5	3165	1.4
134	KNIVTPRTPPPSQ	59000		-0.04	9		32.1	1745	-0.5
0	KGVDAQGTLSKIF	100000			_	10	45.9	340	3.1
		.00000	2001	-0.11	24	15	695.2	2795	0.3

KEY - BB = NUMBER OF THE BACKBONE CHOSEN FROM THE LIBRARY

CLAIMS

- A method for the prediction of the binding affinity of a peptide to a major histocompatibility (MHC) class II
 molecules comprising;
 - a) ascertaining the characteristics of a MHC molecule binding groove,
- b) presenting a selected peptide to the MHC molecule and ascertaining a first conformation score for each pocket bound
 peptide side-chain,
 - c) amending the conformation of each pocket bound peptide side-chain and ascertaining a second conformation score,
 - d) repeating step 3 with alternative conformations of each peptide pocket bound side-chain,
- e) choosing the highest conformation score for each pocket bound peptide side-chain in each binding groove pockets, herein known as 'the pocket', and
 - f) combining the highest conformation score for each pocket and ascertaining a binding score for the complete peptide.
 - 2. A method according to claim 1 which further comprises the step of compiling information on all peptide fragments in a protein and comparing the binding scores.
- 25 3. A method according to any preceding claim wherein the conformation score is ascertained by at least one of the following parameters:
- a) the number of favourable contacts between MHC residues forming one of the pockets and the pocket bound peptide 30 residue; this is value E
 - b) the steric overlap between the pocket bound peptide residue bound in the pocket and an atom forming the pocket; this is value B,
- c) the number of hydrogen bonds which could be formed between 35 the pocket bound peptide residue and an atom forming the pocket; this is value C,
 - d) the strength of electrostatic interactions between any

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polar atoms of the pocket bound peptide residue and any polar atoms forming the pocket; this is value D.

- 4. A method according to claim 3 wherein the steric overlap 5 between the pocket bound peptide residue and the atoms forming the pocket can not be greater than 0.35 Angstroms.
- A method according to claim 3 wherein a favourable contact occurs when an atom from an MHC residue and an atom
 from the peptide residue have their centres separated by no more than the sum of their radii plus 0.5 Angstroms and are not overlapping.
- 6. A method according to the preceding claims wherein values
 15 B to E are imported into a first equation, to give a conformation score (Z)
- 7. A method according to claim 6 wherein the first equation is $Z_n = (cK_2C) (cK_3D) + (cK_4E) (cK_1B)$, where cK_1 to cK_4 are constants and n is the number of the pocket.
 - 8. A method according to claim 7 wherein cK_1 is between 50 and 150.
- 25 9. A method according to claim 7 wherein cK_2 is between 1000 and 2000.
 - 10. A method according to claim 7 wherein cK_{3} is between 250 and 750.
 - 11. A method according to claim 7 wherein cK_4 is between 500 and 1500.
- 12. A method according to any preceding wherein the Z_n value 35 for a pocket is multiplied by a coefficient, L, depending on the pockets importance in binding, to give a second Z_n value.

CLAIMS

- A method for the prediction of the binding affinity of a peptide to a major histocompatibility (MHC) class II
 molecules comprising;
 - a) ascertaining the characteristics of a MHC molecule binding groove,
- b) presenting a selected peptide to the MHC molecule and ascertaining a first conformation score for each pocket bound
 peptide side-chain,
 - c) amending the conformation of each pocket bound peptide side-chain and ascertaining a second conformation score,
 - d) repeating step 3 with alternative conformations of each peptide pocket bound side-chain,
- e) choosing the highest conformation score for each pocket bound peptide side-chain in each binding groove pockets, herein known as 'the pocket', and
 - f) combining the highest conformation score for each pocket and ascertaining a binding score for the complete peptide.
 - 2. A method according to claim 1 which further comprises the step of compiling information on all peptide fragments in a protein and comparing the binding scores.
- 25 3. A method according to any preceding claim wherein the conformation score is ascertained by at least one of the following parameters:
 - a) the number of favourable contacts between MHC residues forming one of the pockets and the pocket bound peptide
- 30 residue; this is value E
 b) the steric overlap between the pocket bound peptide residue
 - bound in the pocket and an atom forming the pocket; this is value B,
- c) the number of hydrogen bonds which could be formed between 35 the pocket bound peptide residue and an atom forming the pocket; this is value C,
 - d) the strength of electrostatic interactions between any

polar atoms of the pocket bound peptide residue and any polar atoms forming the pocket; this is value D.

- 4. A method according to claim 3 wherein the steric overlap 5 between the pocket bound peptide residue and the atoms forming the pocket can not be greater than 0.35 Angstroms.
- A method according to claim 3 wherein a favourable contact occurs when an atom from an MHC residue and an atom
 from the peptide residue have their centres separated by no more than the sum of their radii plus 0.5 Angstroms and are not overlapping.
- 6. A method according to the preceding claims wherein values
 15 B to E are imported into a first equation, to give a conformation score (Z)
- 7. A method according to claim 6 wherein the first equation is $Z_n = (cK_2C) (cK_3D) + (cK_4E) (cK_1B)$, where cK_1 to cK_4 are 20 constants and n is the number of the pocket.
 - 8. A method according to claim 7 wherein cK_1 is between 50 and 150.
- 25 9. A method according to claim 7 wherein cK_2 is between 1000 and 2000.
 - 10. A method according to claim 7 wherein cK_3 is between 250 and 750.
 - 11. A method according to claim 7 wherein cK_4 is between 500 and 1500.
- 12. A method according to any preceding wherein the Z_n value 35 for a pocket is multiplied by a coefficient, L, depending on the pockets importance in binding, to give a second Z_n value.

- 13. A method according to any of the preceding claims wherein all the Z values are summed to give a value J.
- 14. A method according to any of the preceding claims wherein the MHC residue is paired with the pocket-bound peptide residue if an atom from the MHC residue and an atom from the pocket-bound peptide residue have their centres separated by no more than the sum of their van der Waal radii plus one Angstrom.

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- 15. A method according to claim 14 wherein a value A_n is calculated by summing the pairwise interaction frequencies of paired residues.
- 15 16. A method according to either claim 14 or 15 wherein the value A_n for a pocket is multiplied by a coefficient, X, depending on the pockets importance in binding.
- 17. A method according to claim 16 wherein the A_n value for 20 the pockets are summed to give a value P.
 - 18. A method according to any preceding claim wherein the binding score is ascertained by at least one of the following parameters
- 25 a) the number of groove-bound hydrophobic residues; this is value F,
 - b) the number of non groove-bound hydrophilic residues; this is value G,
- c) the number of peptide residues deemed to fit within their 30 respective binding pocket; this is value H.
 - 19. A method according to any one of claims 13 to 18 wherein values F, G, H, J and P are imported into a second equation to give a first binding score, Y.

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20. A method according to claim 19 wherein the second algorithm is $Y=J*F^2*(G*H+1)+P$.

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- 21. A method according to claim 1-17 wherein the hydrophobicity of the pocket bound peptide side chains is evaluated using a hydrophobicity scale; this is value He.
- 5 22. A method according to claim 21 wherein the hydrophobicity scale ranges from -1.8 for lysine to 0.9 for cysteine.
 - 23. A method according to either of claims 21 or 22 wherein $Y=(bK_2C)-(bK_3\ D)+(bK_4E)-(bK_1B)+(bK_5He)+P$.
 - 24. A method according to claim 23 wherein bK_1 is between 1 and 5.
- 25. A method according to claim 23 wherein bK_2 is between 20 15 and 60.
 - 26. A method according to claim 23 wherein bK_3 is between 300 and 900.
- 20 27. A method according to claim 23 wherein bK_4 is between 1 and 20.
 - 28. A method according to claim 23 wherein bK_5 is between 1 and 800.
 - 29. A method according to any preceding claim wherein the steps in claim 3 are repeated for each pocket and each conformation of the peptide residue in said pocket.
- 30 30. A method according to claim 29 wherein the conformation of the peptide is altered by rotating a side chain of the peptide residue by a pre-determined amount.
- 31. A method according to either claim 29 or 30 where in the 35 conformation of the peptide is altered by changing the conformation of the peptide backbone.

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- 32. A method according to any preceding claim wherein the steps are repeated using different peptides from a protein.
- 33. A method according to any of the preceding claim wherein 5 the binding scores (Y) for different peptides are tabulated and compared.
- 34. A method according to any of the preceding claim which is used in the manufacture of a vaccine derived from a peptide10 identified by said method.
- 35. A method according to any of the preceding claims which is used to remove potentially immunogenic sequences from a protein and thus reduce said proteins immunogenicity when administered to an organism.
- 36. A computer conditioned to receive information characterising a peptide bound to the MHC molecule and to utilise said information to perform a procedure having the 20 following steps;
 - a) ascertaining the characteristics of a MHC molecule binding groove;
- b) presenting a selected peptide, which is selected by a predetermined program, to the MHC molecule and ascertaining
 25 a first conformation score;
 - c) amending the conformation of the peptide, by way of a predetermined program, and ascertaining a second conformation score;
 - d) repeating step 3 with other conformations of the peptide;
- 30 e) selecting the peptide conformation with the highest conformation score; and
 - f) calculating the binding score from the conformation score.
- 37. A computer according to claim 36 further comprising a 35 step (7) which comprises repeating steps 1-4 with other peptide fragments in the protein to generate information on all peptide fragments in a protein

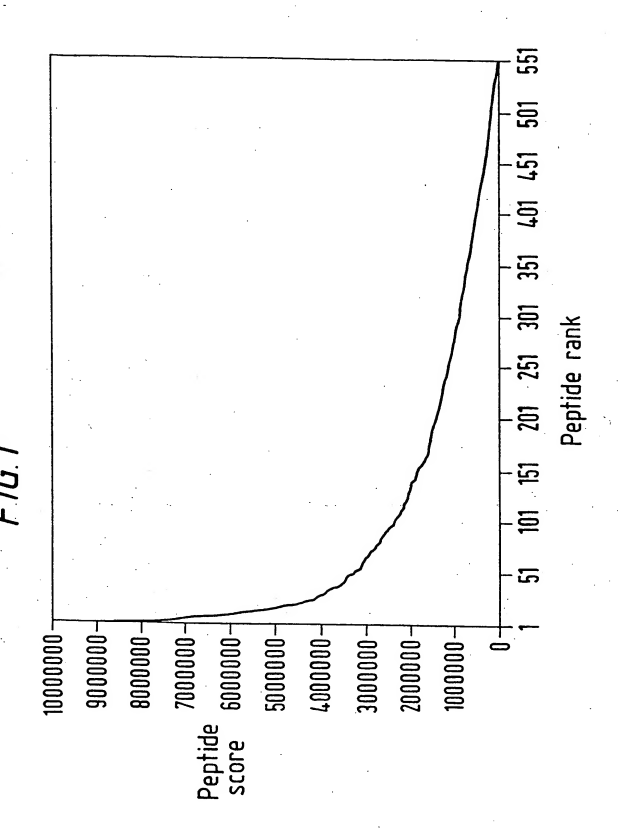
so that a comparison can be made of the strength of the binding between the peptide and the MHC molecule.

- 38. A computer according to either claim 36 or 37 further 5 comprising a step (8) which comprises altering the conformation of the backbone of the peptide fragment.
 - 39. A pharmaceutical composition produced resultant upon to a method as claimed in anyone of claims 1 to 35.

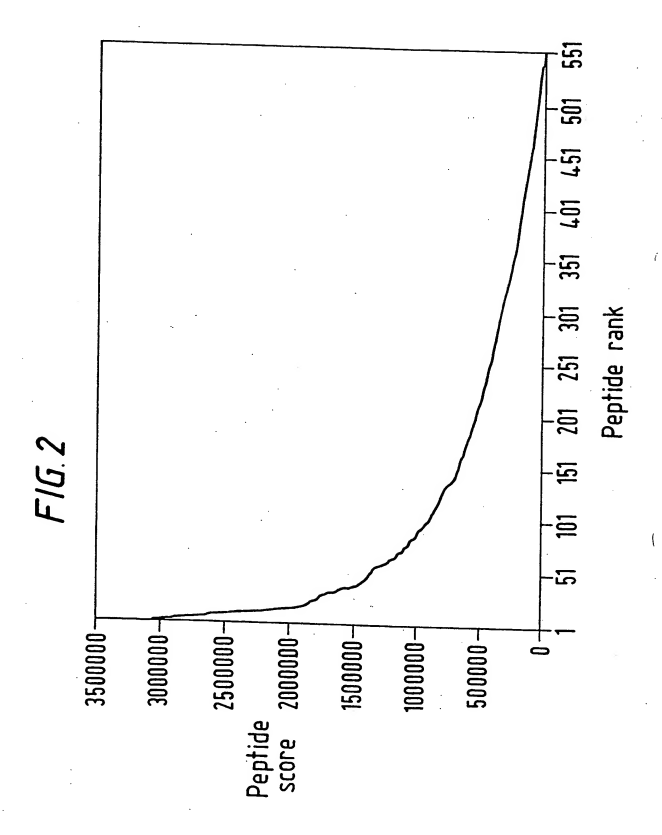
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INTERNATIONAL SEARCH REPORT

International Application No PCT/8/01801

A. CLASSIFICATION OF SUBJECT MATTER
1PC 6 G01N33/569 G01N33/564 G01N33/566 C07K14/705

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols) IPC 6 - 601N - 007K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

Category ·	Citation of decument with indication	
	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Α	WO 95 31483 A (ECLAGEN LTD) 23 November 1995 see page 2, line 23 - line 28 see page 5, line 5 - line 12	1-35
X		39
X,P	WO 97 40852 A (ANERGEN INC) 6 November 1997 see claims 31,32	39
A , P		1-35
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Further documents are listed in the continuation of box C.	χ Patent family members are listed in annex.			
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"A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier document but published on or after the international filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publicationdate of another citation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other means "P" document published prior to the international filing date but later than the priority date claimed	 "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "&" document member of the same patent family 			
Date of the actual completion of theinternational search	Date of mailing of the international search report			
22 October 1998	05/11/1998			
Name and mailing address of the ISA	Authorized officer			
European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040. Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Van Bohemen, C			

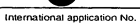
Form PCT/ISA/210 (second sheet) (July 1992)

INTERNATIONAL SEARCH REPORT

International Application No PCT 98/01801

Continu	ation) DOCUMENTS CONSIDERED TO BE RELEVANT		3/01801		
alegory ·	Citation of document, with indication, where appropriate, of the relevant passages	Citation of document, with indication where appropriate, of the relevant passages			
	T.E. JOHANSEN ET AL.: "Peptide binding to MHC class I is determined by individual pockets in the binding groove." SCANDINAVIAN JOURNAL OF IMMUNOLOGY, vol. 46, no. 2, 1 August 1997, pages 137-146, XP002081826 oxford uk see the whole document		Relevant to claim No.		
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## INTERNATIONAL SEARCH REPORT

PCT/GB 98/01801

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)	ヿ
This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:	
Claims Nos.: 36-38 because they relate to subject matter not required to be searched by this Authority, namely:  Rule 39.1(i) PCT - Mathematical method	
Claims Nos.:     because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:	
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).	
Box II Observations where unity of invention is lacking (Continuation of Item 2 of first sheet)	
This International Searching Authority found multiple inventions in this international application, as follows:	
As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.	
As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.	
3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:	
4. No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:	
Remark on Protest  The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.	

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on patent family members

PCT 98/01801

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